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Transcriptional Regulation of *Bacillus subtilis* Glucose Starvation-Inducible Genes: Control of *gsiA* by the ComP-ComA Signal Transduction System

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The Bacillus subtilis glucose starvation-inducible transcription units, gsiA and gsiB, were characterized by DNA sequencing, transcriptional mapping, mutational analysis, and expression in response to changes in environmental conditions. The gsiA operon was shown to consist of two genes, gsiAA and gsiAB, predicted to encode 44.9- and 4.8-kDa polypeptides, respectively. The gsiB locus contains a single cistron which encodes a protein of unusual structure; most of its amino acids are arranged in five highly conserved, tandemly repeated units of 20 amino acids. The 5' ends of gsiA and gsiB mRNAs were located by primer extension analysis; their locations suggest that both are transcribed by RNA polymerase containing sigma A. Expression of both gsiA and gsiB was induced by starvation for glucose or phosphate or by addition of decoyinine, but only gsiA was induced by exhaustion of nutrient broth or by amino acid starvation. Regulation of gsiA expression was shown to be dependent upon the two-component signal transduction system ComP-ComA, which also controls expression of genetic competence genes. Mutations in mecA bypassed the dependency of gsiA expression on Competence genes. Mutations in mecA bypassed the dependency of gsiA expression on Competence genes. Mutations in mecA bypassed the dependency of gsiA expression on Competence genes. Mutations in sporulation but did not otherwise interfere with sporulation, development of competence, motility, or glucose starvation survival. We propose that gsiA and gsii are members of an adaptive pathway of genes whose products are involved in responses to nutrient deprivation other than sporulation.

When available nutrients fall below the levels necessary to sustain rapid vegetative growth of Bacillus subtilis, several developmental programs are initiated. In one program, individual cells undergo a process of morphogenesis that culminates in the differentiation of rod-shaped vegetative cells into spherical, environmentally resistant, dormant cells known as spores. The regulation of this response depends on a number of gene products which appear to be part of a complex, intertwined signal transduction network that controls not only initiation of sporulation but also other programs, such as development of genetic competence, motility and chemotaxis, degradative enzyme synthesis, and antibiotic production. Some of the regulatory genes required for these various adaptive responses are members of the two-component family of bacterial signal transduction systems; others are known to be transcription factors (6, 7, 13). Only limited information exists as to the specific conditions that trigger each response or the order of events as cells enter the stationary phase. It is also not clear whether the responses are mutually exclusive, whether each type of response constitutes an endpoint, or whether at least some of them represent sequential steps in an ordered pathway. It is clear, however, that many gene products induced at the onset of sporulation perform functions that are dispensable for or even inimical to spore formation (46).

Bacterial two-component regulatory systems consist of a histidine protein kinase and a response regulator (47). The histidine kinases, in response to environmental or intracellular information, undergo autophosphorylation and subse-

quently transfer their phosphate groups to their cognate response regulators, in many cases modulating the activity of that protein as a transcriptional activator or repressor. At least four interconnected, two-component regulatory pathways are known to govern transcription of nutrient stress response genes in *B. subtilis*. KinA and Spo0A are the histidine kinase and regulator proteins thought to be responsible for activation of the sporulation pathway (2, 34); ComP and ComA function as the regulatory pair for the competence cascade (6, 7, 49, 51). Other two-component family members in *B. subtilis* are DegU and DegS, which control extracellular enzyme production, and PhoP and PhoR, which control alkaline phosphatase and phosphodiesterase synthesis (for a review, see reference 45).

Much recent interest in B. subtilis has been directed toward elucidation of the environmental cues that initiate development and the search for genes and gene products that mediate it. Several genes that are activated shortly after nutrient deprivation of growing cells have been identified (for reviews, see references 10 and 46). To examine regulation of gene expression during the onset of sporulation in more detail, we have been isolating genes on the basis of their differential expression under conditions of nutrient deprivation (21, 27). This approach involves synthesis of cDNA probes and enrichment for desired sequences by subtractive hybridization. We recently reported the application of this direct method to the identification of two B. subtilis genes rapidly induced in response to glucose deprivation (27). Glucose deprivation is one of the conditions that induce sporulation and several other adaptive responses.

In this report, we characterize the structure and regulation of glucose starvation-inducible genes gsiA and gsiB. Nutri-

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TABLE 1. Bacterial strains and plasmids used in this study

Organism or plasmid	Trait or relevant genotype (plasmid size [kb])	Derivation or reference	
Bacillus subtilis strains			
SMY	Prototrophy	Laboratory stock	
BD1626	hisA1 leuA8 metB5 comA124::pTV55Δ2cat	D. Dubnau; 49	
BD1692	aroD120 mecB31 comA124::pTV55∆2cat comG12::Tn917lac	D. Dubnau; 37	
BD1697	aroD120 mecA42 comA124::pTV55\\Delta2cat comG12::Tn917lac	D. Dubnau; 37	
BD1853	hisA1 leuA8 metB5 comP\(\Delta K1''\)	D. Dubnau; 14	
MB25	Φ(gsiA-lacZ)27 cat	pJPM27→SMY [*]	
MB28	trpC2 pheA1	J. Hoch (JH642); 21	
MB:	ΔamyE::Φ(gsiA-lacZ)42 ermC	pJPM42→SMY	
MB59	gsiB::pJPM22 cat	pJPM22→SMY	
MB60	Φ(gsiB-lacZ)70 cat	pJPM70→SMY	
MB61	ΔgsiA69::cat	pJPM69→SMY	
MB72	gsiA::pJPM20 cat	pJPM20→SMY	
MB82	AgsiB60::neo	pJPM60→SMY	
MB83	AgsiA69::cat AgsiB60::neo	MB61→MB82	
MB84	MB28 AgsiA69::cat	MB61→MB28	
MB186	trpC2 SPβ ^s	H. Taber	
MB223	MB39 comA124::pTV55Δ2cat	BD1626→MB39	
MB225	MB39 comPΔK1	BD1853→MB39	
MB258	MB39 mecA42 comA124::pTV55Δ2cat	BD1697→MB39	
MB259	MB39 mecB31 comA124::pTV55∆2cat	BD1692→MB39	
MB350	MB84 SPβc2del2::Tn917::pSK10Δ6::pJPM117	SPβgsiA+c × MB84	
ZB307A	SPβc2del2::Tn917::pSK10Δ6	P. Zuber; 53	
Escherichia coli strains			
DH5α	F^{-} ϕ 80dlacZ Δ M15 Δ (lacZYA-argF)U169 recA1 endA1 hsdR17 (r_{K}^{-} m $_{K}^{+}$) supE44 λ^{-} thi-1 gyrA relA1	P. Miller	
JM106	JM107 F-	Laboratory stock	
JM107	Δ(lac-proAB) thi gyrA96 endA1 hsdR17 (t _K - m _K -) supE44 (F' traD36 proAB lacPZΔM15) mcrA mcrB	Laboratory stock	
Plasmids			
pAF1	bla cat (11.1)	A. Fouet; 12	
pBEST501/502	bla neo (4.3)	M. Itaya; 17	
pBS-	bla (3.2)	Stratagene, Inc.	
pJPM1	bla cat (3.8)	This work	
pJPM3	bla cat lac Z (8.0)	This work	
pJPM8/9	bla erm (4.5)	This work	
pJPM10/11	bla cat (4.7)	This work	
pJPM15	bla erm (11.6)	This work	
pSGMU2	bla cat (3.7)	J. Errington; 11	
pSGMU32	bla cat lacZ (7.8)	J. Errington; 9	
pSK [±]	bla (3.0)	E. Elliott	
pSP64	bla (3.0)	Promega, Inc.	
pTV20	bla cat erm (15.3)	T. Henkin; 52	

The comPaKI mutation is an in-frame delection that is linked by transformation to a silent kanamycin resistance gene (14).

An arrow indicates construction by transformation.
 SPβ-mediated transduction.

ent exhaustion-induced transcription of gsiA required the products of comP and comA. We propose that gsiA and gsiB are members of an adaptive pathway of genes whose products are involved in nonsporulation responses to nutrient deprivation. In the accompanying report (28), we show that the product of gsiA acts, directly or indirectly, as a negative regulator of the sporulation pathway.

MATERIALS AND METHODS

Bacterial strains and plasmids. The bacterial strains and plasmids used in this work are listed in Table 1. Strains MB258 (mecA42) and MB259 (mecB31) were constructed by congression in the following fashion. Chromosomal DNAs isolated from mecA42 and mecB31 mutant strains (BD1692 and BD1697), which also carried the comA124::pTV55\(\Delta\)cat and comG12::Tn917-lacZ mutations, were used to transform

strain MB39 (gsiA-lacZ), with selection for Cm^r (i.e., for introduction of the comA mutation). Since gsiA-lacZ is entirely dependent on comA for its expression (see Results), the transformants were then scored for expression of \(\beta_{\text{-ga}} \) lactosidase on nutrient broth sporulation medium (DS) (42) plates containing 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (120 µg/ml). For the cross with mecB31 DNA, transformants were also screened for the Spo- phenotype exhibited by mecB mutants (37). To confirm that the comA mec strains did not possess the comG12-lacZ fusion, the gsiA fusion was reintroduced into strain SMY by transformation with selection for Erm'. Erm' transformants were scored for \(\beta\)-galactosidase production on DS plates containing 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside and screened for the Amy phenotype. Both the gsiA and comG fusions carry the emC gene from Tn917, but only the gsiA fusion is expressed in complex medium and is integrated

within the amyE locus. All of the Erm' transformants generated with DNAs from strains MB258 and MB259 exhibited a LacZ⁺ Amy⁻ phenotype on DS plates.

The plasmids used in this study were the following. Integration plasmid pJPM1 was constructed by substituting the 322-bp PvuII fragment of pSGMU2 (11) with the 382-bp PvuII fragment of plasmid pBS (Stratagene, Inc.). Plasmid pJPM1 and its derivatives are unable to replicate in B. subtilis but, upon integration into the chromosome, confer selectable chloramphenicol resistance. pJPM: so contains T3-T7 promoter-primer sequences for plasmid sequencing and in vitro synthesis of uniformly labeled RNA probes. Integrative lacZ transcriptional fusion vector pJPM3, which contains a single BamHI site in the polylinker upstream of lacZ, was constructed by subcloning the 5.0-kb BamHI-Bg/II lacZ-cat cassette from pSGMU32 (9) into the unique BamHI site of pSPo4 (Promega, Inc.). Plasmids pJPM8 and pJPM9 are derivatives of pSK- (Stratagene, Inc.) that carry the erythromycin resistance (Erm^r) gene from plasmid pTV20 (52) in the EcoRV site in opposite orientations. pJPM11 and pJPM12 carry the cat gene from pC194 cloned in the EcoRV site of plasmid pSK⁻ in opposite orientations. Plasmid pJPM15 is an Erm' derivative of pAF1 (12). It is a single-copy integrative vector which allows construction of transcriptional fusions to lacZ and insertion into the amyE gene. Integration of the insert in pJPM15 occurs by double crossover into the chromosome within the α-amylase gene, whose inactivation was confirmed by absence of a halo of starch hydrolysis on TBAB (Difco Laboratories)-starch plates stained with a solution of 1.0% (wt/vol) iodine. Plasmids pJPM67 and pJPM74 were rescued from the chromosome by using conventional cloning techniques (52) following recombinational integration of gsiA- or gsiB-containing plasmids derived from pJPM1.

T3-T7 transcription plasmids. The following plasmids are derivatives of T3-T7 transcription vector pBS⁻ or pSK⁻ (Stratagene, Inc.). Plasmid pCM213, containing the dciA promoter region, was described by Slack et al. (44). pJLB10, a derivative of pSK⁻, contains the gsiA promoter region as a 0.65-kb Pst1-Acc1 fragment (Fig. 1A). Plasmid pJPM17 contains the gsiB promoter region in pSK⁻, as shown schematically in Fig. 1B and 2. pJLB4 contains the spo0H 5' region in pSK⁻ as a 500-bp HindIII fragment from pJOH25 (16). Plasmid pJLB7, which contains the veg promoter region, was constructed by subcloning the 334-bp EcoRI-BamHI fragment of pPH9 (19) into the compatible sites of pSK⁻. pBScitB is a derivative of pBS⁻ which was created by subcloning the 350-bp EcoRI-HindIII fragment from

plasmid pMR41 (38).

General methods. B. subtilis cells were made competent and transformed by the method of Piggot et al. (35). Selection for drug resistance was on DS medium plates containing chloramphenicol (2.5 µg/ml), neomycin (5.0 µg/ml), or erythromycin and lincomycin (0.5 and 12.5 µg/ml, respectively). Strains of Escherichia coli were made competent and transformed by the method of Hanahan (15). Plasmid DNA was isolated from E. coli by a method based on the alkaline lysis method of Birnboim and Doly (1). Restriction endonucleas. and DNA modification enzymes were obtained from New England BioLabs, Inc., and used as recommended by the supplier.

Growth and sporulation conditions. B. subtilis cells were induced to sporulate either by medium replacement (resuspension) or by nutrient exhaustion. For resuspension experiments, cells in the exponential growth phase (~100 Klett units) in medium 121J (Tris-buffered glucose-glutamate me-

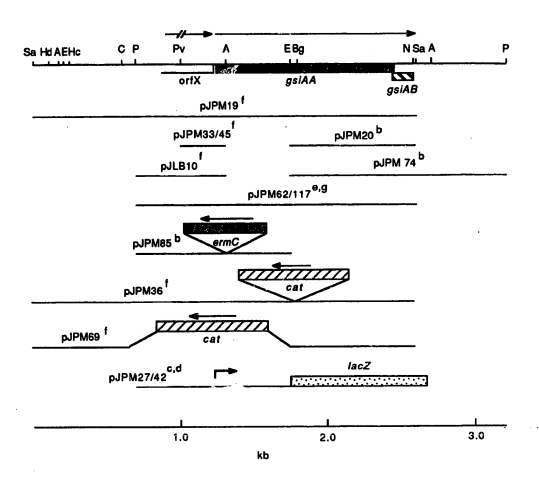
dium; 27) were harvested and resuspended in medium that lacked glucose, NH₄Cl, or K₂HPO₄ for carbon, nitrogen, or phosphate starvation, respectively. Medium 121CG differed from 121J in that glucose was replaced with 1.0% Casamino Acids. Medium 121F was prepared as described by Slack et al. (44). Time zero (T_0) is defined as the time of resuspension in starvation medium. For exhaustion experiments, an exponential-phase culture in DS medium was diluted to give an A_{600} of 0.05. Culture growth at 37°C was monitored by measuring A_{600} ; T_0 was defined as the end of exponential growth. Induction of sporulation with decoyinine has already been described (44). The number of heat-resistant spores was determined after 12 to 24 h at 37°C. Samples were plated either before or after heat treatment (80°C, 20 min) to measure the number of viable or heat-resistant CFU, respectively.

RNA isolation and RNase protection analysis. Isolation of total RNA from B. subtilis was done by the guanidinium isothiocy anate method (44). Cells were harvested during exponential growth in 121J medium and 1 h after starvation for glucose (T_1) or at 5-min intervals after addition of decoyinine (500 μ g/ml) to 121F medium. Antisense RNA probes were synthesized by using the Stratagene riboprobe system. RNA hybrids were treated with RNase T2 as described by Slack et al. (44). Nuclease-resistant hybrids were separated by electrophoresis in 5% polyacrylamide-7 M urea sequencing gels and visualized by autoradiography.

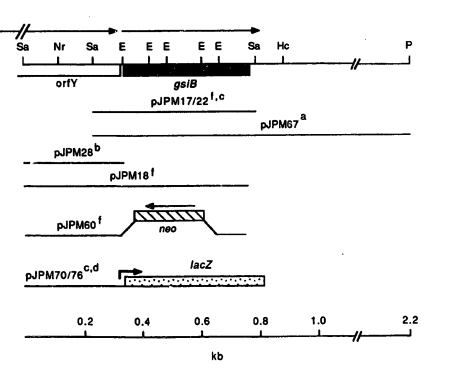
Identification of the gsiA and gsiB transcription start sites. Primer extension reactions were performed as described by Sambrook et al. (40). The primers were synthetic 19-nucleotide DNA oligonucleotides complementary to the 5' terminal region of gsiA or gsiB mRNA (Fig. 3). Sixty picomoles of oligomer was end labeled by incubation with 150 µCi of ²P]ATP (5,000 Ci/mmol) and T4 polynucleotide kinase as described previously (40). Free nucleotides were separated from labeled oligonucleotide by three precipitations with ammonium acetate-ethanol. For primer extension reactions, 6 ng of oligomer (5 \times 10⁴ cpm) was annealed with 20 μg of RNA in a buffer containing 80% formamide, 0.5 M NaCl, 1 mM EDTA, and 40 mM piperazine-N,N'-bis(2-ethanesulfonic acid) (PIPES; pH 6.8). The mixture was incubated at 80°C for 3 min and then cooled gradually to 30°C. The hybridized nucleic acids were precipitated with ethanol, dissolved in 20 µl of reverse transcriptase buffer (50 mM KCl; 1 mM EDTA; 55 mM Tris-HCl [pH 8.0]; 5 mM dithiothreitol; 8 mM MgCl₂; dATP, TTP, dGTP, and dCTP each at 2.5 mM) and incubated with 20 U of avian myeloblastosis virus reverse transcriptase (Life Sciences; St. Petersburg, Fla.) at 42°C for 45 min. Four microliters of a solution containing 0.25 M EDTA and 10 µg of RNase A per ml was added, and the incubation was continued at 37°C for 30 min. The reaction products were extracted with phenolchloroform, precipitated with ethanol, solubilized in 5 µl of loading buffer, and separated by electrophoresis in a 5% polyacrylamide-7 M urea gel.

DNA sequencing. Fragments of DNA from plasmids pJPM18 and pJPM19 were cloned into phagemids pSK⁺ and pSK⁻ for sequencing by the dideoxy-chain termination method of Sanger et al. (41) with modified T7 DNA polymerase (Sequenase, Version 2.0; U.S. Biochemical Corp.). When necessary, reactions using *Taq* DNA polymerase (Cetus) and dGTP analogs were used to resolve sequence compressions. Single-stranded DNA templates were prepared from R408 helper phage lysates of superinfected *E. coli* JM107 cells harboring recombinant pSK plasmids, as recommended by Russel et al. (39). The nucleotide sequence

Α.



B.



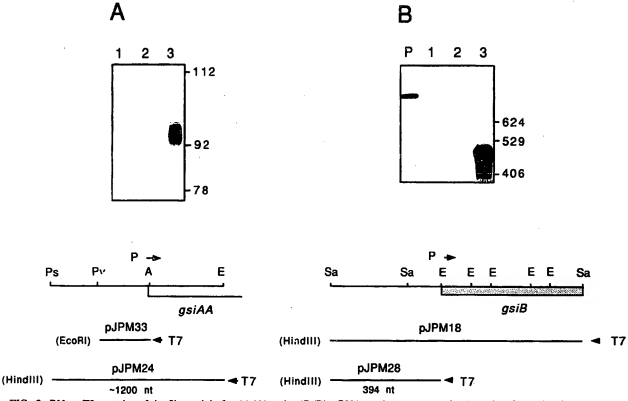


FIG. 2. RNase T2 mapping of the 5' termini of gsiA (A) and gsiB (B) mRNAs and appearance of gsiA and gsiB mRNAs in response to glucose deprivation. The 351-bp PvuII-AccI gsiA fragment and the 521-bp Sau3A gsiB fragment were used for synthesis, in the direction indicated, of uniformly labelled antisense RNA probes specific for each sequence. Labeled gsi probes were separately hybridized with 20- μ g samples of total B. subtilis RNA from cells collected at T_0 in 121J medium (lane 2) or T_1 in 121J lacking glucose (lane 3) or with RNA from Saccharomyces cerevisiae (lane 1). P, untreated probe (5,000 cpm). The hybridization products were treated with RNase T2, and the nuclease-resistant products were resolved on a 5% polyacrylamide-7 M urea sequencing gel. The samples were run alongside end-labeled Hpall fragments of pBR322, and the sizes are shown in base pairs on the right. nt, nucleotides. For restriction site abbreviations, see the legend to Fig. 1. Ps, PstI.

of both strands was determined by using a series of synthetic oligonucleotides that prime at intervals of approximately 350 nucleotides. Oligonucleotides were synthesized by using an Applied Biosystems 380B DNA synthesizer at the Tufts Protein Chemistry Facility. Sequence analysis was performed by using the University of Wisconsin Genetics Computer Group package (4).

Construction of gsiA-lacZ and gsiB-lacZ transcriptional fusions. Plasmid pJPM27 is a gsiA-lacZ transcriptional fusion. It was constructed by cloning the 1.2-kb PstI-Bg/II fragment carrying the gsiA promoter in front of the lacZ gene of vector pJPM3 (Fig. 1A). Plasmid pJPM70 is a gsiB-lacZ transcriptional fusion constructed by cloning the 0.36-kb Sau3A-EcoRI fragment carrying the gsiB promoter (purified from a multiple cloning site as a HindIII-BamHI fragment) into HindIII-BamHI-digested pJPM3 (Fig. 1B). The gsiA-lacZ and gsiB-lacZ fusions were inserted into the chromo-

some by single-reciprocal recombination by transformation of competent cells of strain SMY with plasmids pJPM27 and pJPM70 and selection for chloramphenicol resistance (Cm'). Chromosomal DNA from one chloramphenicol-resistant transformant was purified from each cross and subjected to Southern hybridization analysis to confirm that pJPM27 and pJPM70 had integrated in a single copy at the chromosomal gsiA and gsiB loci, respectively (data not shown).

Plasmid pJPM42 was constructed by subcloning the 3.2-kb HindIII-SacI fragment of pJPM27 into the HindIII-SacI backbone of pJPM15. pJPM42 was linearized with NruI and used to transform SMY to Erm^r. Since the constructs carried homology to the amyE locus, this transformation resulted in integration of the fusion constructs by replacement recombination at this locus.

Deletion of the chromosomal gsiA and gsiB genes. Plasmids containing in vitro-derived deletions and insertions of gsiA

FIG. 1. Cloned DNA from the gsiA (A) and gsiB (B) regions of the B. subtilis chromosome. Physical maps of the DNA inserts in the plasmids used are shown below the abbreviated restriction maps. The locations of the gsi genes are indicated. The positions and structures of various insertions in the chromosome are also indicated. Restriction sites: E, EcoRI; Hd, HindiII; Pv, PvuII; B, BgiII; A, AccI; Hc, HincII; C, ClaI; N, NruI; Nr, NarI; Sa, Sau3A; P, PstI. Vectors are indicated by the following superscripts: a, pBS; b, pJPM1; c, pJPM3; d, pJPM15; f, pSK⁻; g, pBEST501.

A.

Q L G E L Y V S S A L Q W H Q E E N H H I E T M V K D M T G E Q 101 >>>>>>>>>> Q R G S W M E R N K V G F H *

11 qsiAA олиз MKQTIPSSYVGLKINEWYTHIRQFHVAEAERVKL 401 EVEREIEDMEEDQDLLLYYSLMEFRHRVMLDYI 501 P F G E D T S Q L E F S E L L E D I E G N Q Y K L T G L L E Y N F F R G M Y E F K Q K M F V S A M M Y Y K R A E K N L A L V S D 701 D I E K A E F A F K M A E I F Y N L K Q T Y V S M T Y A V Q A L AACATACCAAATGTATGAAACGTACACCGTCCGCAGAATCCAATGTGAATTCGTTATTGCAGGTAATTATGATGATATGCAGTATCCAGAAAGAGCATTG D D M Q Y P E R A TYQMYETYTVRRIQCEFVIAGNY $\tt CCCCACTTAGAACTGGCTTTAGATCTTGCAAAGAAGAAGGCAATCCCCGCCTGATCAGTTCTGCCCTATATAATCTCGGAAACTGCTATGAGAAAATGG$ H L E L A L D L A K K E G N P R L I S S A L Y N L G N C Y E K M G ELQKAAEYFGKSVSICKSEKPDNLPHSIYSLTQ 1101 AGTTCTGTATAAACAAAAAAATGACGCCGAAGCGCAAAAAAAGTATCGTGAAGGATTGGAAATCGCCCGTCAATACAGTGATGAATTATTTGTGGAGCTT V L Y K Q L N D A E A Q K K Y A E G L E I A R Q Y S D E L F V E L Q F L H A L Y G L N I D T E S V S H T F Q F L E E H M L Y P Y I E ELAHDAAQFYIENGQPEKALSFYEKMVHAQKQI 1401 CCAGAGAGGAGATTGTTTATATGAAATCTAAATGGATGTCAGGTTTGTTGCTCGTTGCGGGTTCAGCTTTACTCAGGTGATGGTTCATGCAGGTGA Q R G D C L Y E I *
gsiAB M K S K W M S G L L L V A V G F S F T Q V M V H A G E

1501 AACAGCAAACAGAAGGGAAAACATTTCATATTGCGGCACGCAATCAAACATGATGCATAAAAAAAGACCCTTAGGGGTCTTTTTTATTTCTTCAGCTT T A N T E G K T F H I A A R N Q T * >>>>>>>

1601 CCATTCTTTTATCGTCAGCTCAGAAGATC

В.

GATCAAGACCGTACATATAAGAATGTCGCTTCTCAAATCCAAGGCTGGCGAGAAGTCGTTTTGGGCTATCGAGACACGTTTGGCTGGAAAAAACTTTTCC D Q D R T Y K N V A S Q I Q G W R E V V L G Y R D T F G W K K L F AGATACTGCCGGTTGCCGGAATGGTTTTTGGCGCCGCTGCCAATCGCTCAACATTAAACGACATTACCGAGACAGGCATGATGCTGTACAAAAAAGAGGCG I V P V A G M V F G A A A N R S T L N D I T E T G M M L Y K K R R CATTCTTGAACGACTGAAAGAACAGAACGAGAGATGGAATAGCAGAAAGCAGACGGACACCGCGATCCGCCTGCTTTTTTTAGTGGAAACATACCCAAT <<<< <<<<

ILERLKETEREME >>>>>>>

OJM1 301 GTGTTTTGTTTGTTTAAAAGAATTGTGAGCGGGAA<u>TACAAC</u>AACCAACACCAACTT<u>AAAGGAGG</u>AATTCAAAATGGCAGACAATAACAAAATGA<u>GCAGAGA</u>

gsib m a d n n l m s r e <u>AGAAGCAGGTAG</u>AAAAGGCGGAGAAACAACAAGCAAGAACCATGACAAAGAATTCTATCAAGAGATTGGTCAAAAAGGCGGAGAAGCCACTAGCAAAAAAC

E A G R K G G E T T S K N H D K E F Y O I G O K G 501

N H D K E F Y O E I G E GAGAAGCCACTAGCGAAAATCATGACAAAGAATTCTATCAAGAAATCGCCGAAAAGGCCGAGAGGCAACAAGCAAAAATCATGATAAAGAATTCTACCA

D K E F Y O E I G R K G G E A T S K N H D K <u>I</u> G S K G G N A R N N D *

801 <<<<<<<

1001 CACCATCACCCTGACTCAGACG

FIG. 3. DNA sequences of the nontemplate strands of gsiA (A) and gsiB (B). The derived amino acid sequences of the gsi genes are shown below the nucleotide sequences. Single underlining identifies the nucleotide sequences complementary to oligonucleotides OJM3 and OJM1, which were used for primer extension mapping. The vertical arrows identify the proposed 5' ends of the gsi transcripts. The presumed -10 and -35 regions are shown underlined. The putative ribosome-binding sites are doubly underlined. The proposed ComA box upstream of the gsiA promoter is shown in boldface. The stems of putative transcription terminator sequences are shown as converging arrowheads. The 20-amino-acid repeats of the proposed GsiB protein are shown by single underlinings.

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and gsiB (Fig. 1A and B) were linearized and used to transform competent cells of strain SMY to antibiotic resistance. Mutations of gsiA and gsiB constructed by gene replacement were confirmed by Southern blot analysis (data not shown).

Insertion of gsiA at the SPB locus. Plasmid pJPM117 (Fig. 1A) carries the complete gsiA operon and all of the control sequences of the gsiA promoter. This plasmid was constructed by ligating the 2.0-kb fragment of pJPM19 into the Hindli and Pstl sites of pBEST501 (17). Plactid pJPM117 was used to transform competent cells of strain ZB307A to neomycin resistance (Neo') (32). The Neo' transformants were pooled, and an SPB transducing lysate was generated that contained a bacteriophage mixture, some of which carried pJPM117. To obtain a pure SPβgsiA⁺ lysogen, the lysate was used to infect exponential-phase cells of SPBsensitive strain MB186 with selection for neomycin resistance (53). A Neo' iysogen of MB186 was used as a source of an SPBgsiA+ specialized transducing phage.

Measurement of β-galactosidase activity. β-Galactosidase

was assayed as previously described (22, 44).

Nucleotide sequence accession numbers. The DNA sequences presented in this report have been forwarded to the EMBL, GenBank, and DDJL nucleotide sequences data libraries under accession numbers X56679 (gsiA) and X56680 (gsiB).

RESULTS

Analysis of gsiA and gsiB transcription units. We previously described the isolation of DNA segments designated gsiA and gsiB from a \(\lambda ZAP\) library of B. subtilis chromosomal DNA by use of cDNA probes enriched for glucose starvation-induced transcripts (27). To define the relevant transcription units more precisely, total RNA was prepared from cells growing exponentially in a medium containing 0.5% glucose and from cells harvested 1.0 h after resuspension in the same medium lacking glucose. The RNA was annealed to uniformly labeled RNA probes spanning the cloned gsiA and gsiB regions in both orientations, and hybrid formation was tested by treatment with RNase T2. The direction of transcription of gsiA and gsiB was judged by the pattern of protection.

Previous studies using cDNA probes prepared from glucose-starved cells localized the gsiA transcription unit to the 1.6-kb PvuII-Sau3A fragment carried on plasmid pJPM24 (27; Fig. 1 and 2A). About 635 bases of an RNA probe transcribed from this fragment were protected following hybridization to RNA from glucose-starved cells (data not shown). This established an approximate location of the 5' end of the gsiA transcriptional unit within the 0.35-kb PvuII-AccI fragment. The same RNA protected a segment of an antisense probe, produced by using pJPM33, that corresponded approximately to the 94 bases to the left of the AccI site shown in Fig. 2A. RNA from glucose-starved cells protected approximately 440 bases of a gsiB probe made from plasmid pJPM18, suggesting that the 5' end of the gsiB transcriptional unit lies near the leftmost EcoRI site (Fig. 2B). We failed to detect any signal after hybridization of labeled antisense RNA probes prepared from plasmid pJPM28. The precise location of the 5' end of gsiB mRNA was resolved by nucleotide sequence analysis and primer extension experiments (see below). The gsiA and gsiB transcripts were not detectable in exponentially growing cells, and an increase in the steady-state levels of gsiA and

gsiB transcripts was observed within 1.0 h after resuspension (Fig. 2A and B).

DNA sequence analysis of gsiA and gsiB. Figure 1 shows a genetic and physical map of the gsiA and gsiB regions of the B. subtilis chromosome. The nucleotide sequences of the 1.6-kb PvuII-Sau3A fragment containing gsiA and a 1.0-kb region of the gsiB locus were determined on both strands (Fig. 1 and 3). The sequenced region of gsiB extends from the upstream Sau3A restriction site to 236 bp downstream of the downstream Sau3A restriction site (Fig. 1 and 3B). The sequence of the gsiA locus revealed two open reading frames, designated gsiAA and gsiAB, which are capable of encoding proteins of 378 and 44 residues, with deduced molecular weights of 44,959 and 4,793, respectively. gsiAA and gsiAB overlap by eight nucleotides and may be transcribed as an operon (Fig. 3A). The putative ATG start codons of gsiAA and gsiAB are preceded by sequences (indicated in Fig. 3) that should, in principle, serve as ribosome-binding sites. A translational fusion of gsiAA to lacZ encoded B-galactosidase activity, indicating that the gsiAA reading frame is used in vivo (data not shown). It is not known whether gsiAB actually encodes a protein.

DNA sequence analysis of the g B locus revealed a single open reading frame which could code for a protein of 123 amino acids with a calculated molecular weight of 13,789. The putative gsiB protein has an unusual structure consisting of multiple direct repeats organized in five tandem units of 20 amino acids (Fig. 3B). The putative start codon of gsiB is preceded by an apparent ribosome-binding site, and a translational fusion of lacZ to gsiB was active, indicating that this open reading frame is used in vivo (data not shown). The primary products of the gsiA operon and the gsiB gene are

predicted to be markedly hydrophilic.

DNA sequence analysis also showed the presence of partial protein-coding sequences upstream of both gsiA and gsiB (Fig. 3) which had no significant homology with proteins in the GenBank data base. Furthermore, inactivation of orfX and orfY did not affect growth or sporulation.

Immediately downstream of the gsiAB and gsiB stop codons are sequences which show strong similarity to factorindependent transcription termination sites (i.e., sequences capable of coding for an RNA stem-loop structure followed by a uridine-rich sequence). The location of these sites is consistent with the apparent locations of the 3' ends of gsiA and gsiB transcripts as determined by nuclease protection

experiments (data not shown).

Mapping of the 5' termini of gsiA and gsiB mRNAs. The results of low-resolution experiments suggested a single region of initiation for each of the transcription units and, in conjunction with DNA sequence analysis, allowed the design of chemically synthesized DNA oligonucleotides which were used to identify the mRNA start sites for gsiA and gsiB transcripts induced by glucose starvation. The glucose starvation-induced promoters for gsiA and gsiB were mapped to a region immediately upstream from GsiAA and GsiB by extension of 19-nucleotide-long synthetic primers called OJM3 and OJM1, respectively (Fig. 3). Each primer was labeled at its 5' end, annealed to total RNA isolated from cells collected after 60 min of glucose starvation, and extended by using reverse transcriptase. The primer extension products were then electrophoresed together with dideoxynucleotide chain termination reactions carried out by using the same primer (Fig. 3A and B). The 5' termini of gsiA and gsiB mRNAs were 37 and 21 bases upstream of their respective putative translation initiation codons (Fig. 3 and 4). No primer extension products were obtained with RNA

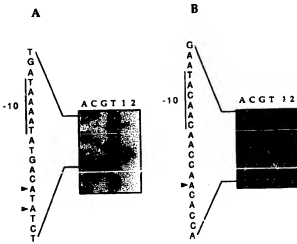


FIG. 4. Determination of the 5' termini of the gsiA (A) and gsiB (B) transcripts by primer extension. A 32P-end-labeled oligonucleotide primer was annealed to 20 µg of total cellular RNA and extended by using deoxynucleoside triphosphates and reverse transcriptase. The DNA products were separated by electrophoresis on a 5% polyacrylamide-7 M urea gel. Their mobilities were compared with a dideoxynucleotide sequencing ladder (lanes A, C, G, and T) produced with the identical primer and the sense strand of singlestranded pJPM45 (gsiA) or pJPM17 (gsiB) as the template. Strain SMY was grown in 121J medium, and RNAs were extracted from exponentially growing cells (lane 2) and at 60 min after glucose starvation (lane 1). The complement of the sequence surrounding the inferred transcription start point is indicated. The complements of the nucleotides that correspond to the start sites of gsiA and gsiB mRNAs are indicated by arrowheads.

from exponentially growing cells (Fig. 4). These findings are in accordance with the transcription start points defined by RNase protection experiments.

Figure 3 shows that the sequences upstream of the putative start sites of gsiA and gsiB mRNAs have limited homology to promoters recognized by the $E\sigma^A$ form of B. subtilis RNA polymerase. The -10 sequence deviated from the consensus TATAAT by one nucleotide for gsiA and by two nucleotides for gsiB. Located upstream from these possible -10 sequences are rather poor -35 sequences. Promoters that require positive regulators often have poor -35 sequences (36). We have been unable to detect transcription of gsiA or gsiB in vitro with purified EoA RNA polymerase (29). The gsiA and gsiB promoter regions contain no matches to the consensus sequences for alternate forms of B. subtilis RNA polymerase (3, 24).

gsiA and gsiB promoters are responsive to various forms of nutrient limitation. We monitored the transcriptional responses of the gsiA and gsiB genes during growth and sporulation by using gsiA- and gsiB-lacZ gene fusions. Fusion plasmids pJPM27 and pJPM70 (Fig. 1) were integrated into the chromosome of wild-type strain SMY, and the levels of β-galactosidase in response to various environmental conditions were measured. As expected, both genes were expressed at a low level during exponential growth in medium containing excess glucose and were induced rapidly in response to glucose limitation (Fig. 5A and D). This effect was reversed immediately by restoration of glucose to starved cells (data not shown). Expression of gsiA and gsiB was also assessed under other conditions that lead to efficient sporulation. Both promoters (in particular, gsiA) were

induced by decoyinine (Fig. 5B and E), a purine analog that inhibits GMP synthetase (20, 23). The gsiA-lacZ fusion was also turned on, albeit transiently, when cells in DS medium entered the stationary phase (Fig. 5C), but under similar conditions, gsiB-directed β-galactosidase activity was no higher than the background level of the endogenous β-galactosidase activity normally present in B. subtilis (Fig. 5F).

To assess whether the gsiA and gsiB promoters were generally responsive to other forms of nutrient limitation, we examined their activities in media deficient in a single primary nutrient. The results of these experiments indicated that gsiA-lacZ was induced under all conditions of nutrient limitation, while the gsiB-lacZ fusion was induced by phosphate limitation but unaffected by nitrogen or amino acid

starvation (Table 2).

Appearance of gsiA and gsiB transcripts in response to decoyinine. The rates of response of gsiA, gsiB, and other genes to decoyinine treatment were estimated by hybridization experiments. As shown in Fig. 6A and B, levels of gsiA and gsiB steady-state RNAs increased within 5 min after treatment with decoyinine. By comparison, the steady-state level of the dciA transcript (which encodes a dipeptide transport system) increased between 5 and 10 min after drug addition (Fig. 6A), as previously reported (21, 44). Changes in the steady-state levels of citB (aconitase) and spo0H (σ^H) RNAs were generally consistent with previous results (5, 16). The steady-state level of veg (a vegetative cell transcript with an unknown function) was essentially unaffected by treatment with decoyinine (Fig. 6), as expected from previous work (12a). The changes we observed reflect a very rapid and significant change in the abundance of gsiA and gsiB mRNAs in response to the environmental signal.

The gsiA operon is controlled by the ComP-ComA signal transduction system. Results from several laboratories are consistent with the hypothesis that the B. subtilis twocomponent sensor-regulator system comP-comA (49, 51) responds to the availability of the carbon and nitrogen source or to a combination of these signals. comA and comP are required for competence gene expression, surfactin production, and transcription of at least one positive regulator of degradative enzyme synthesis (26, 30, 32, 49, 51). Since gsiA transcription was strongly induced in response to glucose and nitrogen deprivation and since gsiA mutations confer a catabolite-resistant sporulation phenotype (see below), we investigated the dependence of gsiA expression on comP and comA. Strains MB223 (comA124::pTV55Δ2) and MB225 (comPΔK1), each of which carries a gsiA-lacZ transcriptional fusion at the chromosomal amyE locus, showed a drastic reduction in \(\beta \)-galactosidase activity in DS medium compared with the com⁺ parent (MB39) (Fig. 7). The β-galactosidase activity directed by the gsiA promoter in strain MB39 increased sharply after the beginning of the stationary phase with the same induction ratio and kinetics as the corresponding fusion integrated at its normal locus, indicating that all of the cis-acting sequences required for gsiA transcription were carried by the PstI-BglII fragment. Since gsiB is not expressed in DS medium, its dependence on ComA-ComP was not tested. The dependence of gsiA expression on other regulatory genes known to affect competence was also tested (6, 7, 14, 37, 50). Induction of gsiA-driven β-galactosidase activity in response to nutrient exhaustion was independent of spo0A and abrB (see the accompanying report [28]), srfA, comQ, spo0KA(oppA), spo0KE(oppE), degUS, sin, and spo0H (data not shown).

The loci mecA and mecB are defined by mutations that suppress the competence defect of comA mutants and allow

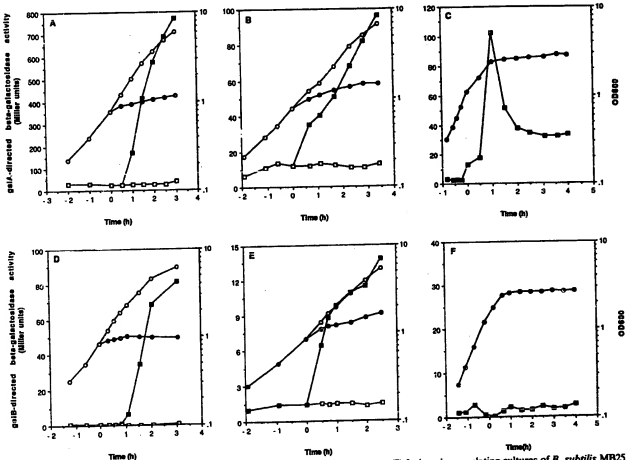


FIG. 5. Expression of β -galactosidase from gsiA-lacZ (A to C) and gsiB-lacZ (D to F) fusions in sporulating cultures of B. subtilis MB25 and MB60, respectively. Samples were assayed at the indicated times for growth (circles) and β -galactosidase activity (squares). Panels: A and D, 121J medium (open symbols) and 121J medium from which glucose was removed at T_0 (closed symbols); B and E, 121J medium (open symbols) and 121J medium to which decoyinine was added to 500 µg/ml at T_0 (closed symbols); C and F, DS medium (closed symbols). The abscissa is divided into hours preceding or following the end of exponential growth, which is indicated as time zero.

otherwise wild-type cells to express competence genes under inappropriate nutritional conditions (8, 37). Mutations in mecA suppressed the ComA dependence of gsiA expression and restored normal growth stage-specific regulation (Fig. 7). In a comA mecB mutant strain (MB259), gsiA-lacZ expression was partially restored (maximum activity, 15 U), compared with comA strain MB223 (1 to 2 U) (data not shown).

Gene inactivation of the gsiA and gsiB loci. We constructed deletion-insertion mutations in the cloned gsiA and gsiB genes in vitro (Fig. 1), which were then used to replace the wild-type alleles in the chromosome of strain SMY with a gene that confers antibiotic resistance by homologous recombination. Chromosomal deletion of gsiA or gsiB or both yielded no obvious effects on bacterial growth or viability in complex or minimal media containing various fermentable or nonfermentable carbon substrates. We saw no significant differences in the abilities of the mutant cultures to form spores (Table 3 and data not shown), establish competency, develop motility, or survive glucose starvation. The lack of a phenotype associated with deletion of gsiA or gsiB or both

might result from the existence of other genes with similar or overlapping functions.

Because both gsiA and gsiB are induced by glucose deprivation, we considered a role for these genes in catabolite repression. We discovered that gsiA (but not gsiB) mutants have a defect in catabolite repression of sporulation (Crs- phenotype). Under conditions which normally repress sporulation (2.0% glucose), a gsiA mutant was extremely efficient in the formation of heat-resistant endospores (Table 3). To demonstrate that the Crs- phenotype did not result from a polar effect of the chloramphenicol resistance gene, gsiA+ was restored to the AgsiA strain on a specialized tranducing phage. As shown in Table 3, the Crs phenotype was abolished when a wild-type copy of the gsiA operon was provided in trans at the SPB locus. Disruption of gsiA did not relieve glucose repression of extracellular protease or amylase production, however, indicating that it does not cause a general release of catabolite repression. Transcription of gsiA was induced in stationary-phase wild-type cells in nutrient broth sporulation medium containing 1% glucose,

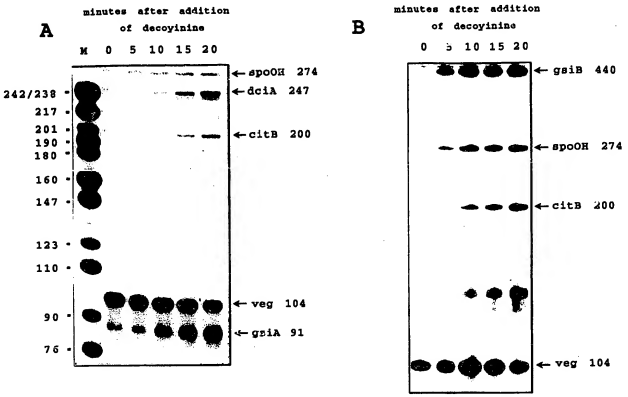


FIG. 6. Time course of appearance of sporulation-associated transcripts in response to decoyinine. Labeled antisense probes were hybridized to 20 µg of B. subtilis RNA purified from SMY cells in 121F medium harvested at the onset (0) and at 5-mi intervals after addition of decoyinine (500 µg/ml). Hybrids were treated with RNase 12 and subjected to electrophoresis in a 5% polyacrylamide-7 M urea gel alongside end-labeled Hpa II fragments of plasmid pBR322 (lane M). The probes were synthesized from pBScitB (citB, 434 nucleotides), pCM213 (dcitA, 738 nucleotides), pJLB4 (spo0H, 589 nucleotides), pJLB10 (gsiA, 713 nucleotides), apply pJLB10 (gsiA, 713 nucleotides), pJLB4 (spo0H, 589 nucleotides), pJLB10 (gsiA, 713 nucleotides), pJLB4 (spo0H, 589 nucleotides), pJLB10 (gsiA, 713 nucleotides), pJLB40 (gsiA, 713 nucleotides), p (gsiB, 634 nucleotides). The numbers in parentheses designate the sizes of the antisense RNA probes. Arrowheads indicate the positions (in bases) of the protected portions of the labeled antisense RNA probes. Lane M contained molecular size markers.

but the time of peak activity was delayed by approximately 1 h (data not shown).

DISCUSSION

The gsiA and gsiB loci of B. subtitis were initially identified and isolated because they are induced by glucose

TABLE 2. Responsiveness of the gsiA and gsiB promoters to various conditions of nutrient limitation

	β-Galactosidase activity (Miller units)				
Limiting nutrient	gsiA-	lacZ	gsiB-lacZ		
	Exponential	Stationary	Exponential	Stationary	
Glucose	8	771	1	- 74	
Nitrogen	5	105	1	1	
Phosphate	6	515	1	190	
Casamino Acids	8	95	1	1	

^a B. subtilis MB25 (gsiA-lacZ) and MB60 (gsiB-lacZ) were grown in 121J or 121CG medium. Exponential-phase samples were collected when the turbidity of the culture gave an Ann of 0.7. Stationary phase samples were collected approximately 3 h after resuspension of cells in medium lacking the indicated nutrient. Stationary-phase activities were normalized by subtracting the activity in control cultures. The activities shown are averages of two experdeprivation (27). The gsiA locus is required for normal catabolite control of sporulation, suggesting that it plays some role in sensing or responding to carbon sources. In fact, gsiA and gsiB are induced by many different changes in nutritional status. They may respond, therefore, to a common signal generated by multiple forms of nutritional deprivation or to multiple signals specific to different stresses.

We have shown that a ComA-dependent mechanism activates gsiA expression in response to an undefined signal associated with exhaustion of nutrient broth. Expression of gsiA in DS medium is transient and seems to represent a special branch of the ComP-ComA pathway, since transcription of gsiA is independent of the products of intermediate and late competence genes (37). ComA also appears to be necessary for induction of gsiA gene expression by other forms of nutrient deprivation (e.g., glucose, nitrogen, and phosphate), as well as by treatment with decoyinine (43). The similarity between various transcriptional activators and the ComA protein and the requirement for ComA in the transcription of gsiA-lacZ raise the possibility that ComA functions directly as a positive regulator of gsiA transcription as cells enter the stationary phase. Indirect mechanisms cannot be ruled out, however (see below).

The regulation and comP-comA dependency of gsiA-lacZ expression resemble those of the degQ gene, which codes for a putative positive regulatory protein for degradative

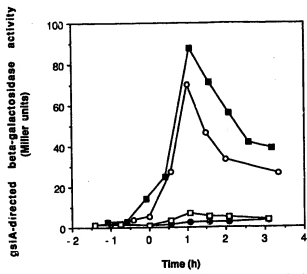


FIG. 7. Dependence of gsiA-directed β-galactosidase synthesis on comA and comP. Production of β-galactosidase in DS medium as a function of growth stage in wild-type, comA124::pTV55Δ2, and $comP\Delta K1$ cells carrying gsiA-lacZ. Symbols: O, MB39 $(com^+\Delta amyE::\Phi gsiA$ -lacZ emC); \Box , MB223 (comA124::pTV55Δ2); \blacksquare , MB225 $(comP\Delta K1)$; \blacksquare , MB258 (comA124::pTV55Δ2) mecA42). The abscissa is divided into hours preceding or following the end of exponential growth, which is indicated as time zero.

enzyme synthesis (25, 26). Another B. subtilis transcription unit whose expression is dependent on comP-comA is srfA, an operon required for production of the antibiotic surfactin. In this case, at least one gene required for establishment of competence is encoded within this large operon (18, 30, 31, 32, 48). Transcription of srfA, degQ, and gsiA is responsive to deprivation of glucose, nitrogen, or phosphate and to decoyinine treatment (18, 25, 26, 32). A proposed target site for ComA-dependent regulation of degQ and srfA was reported to be located in a dyad symmetry element just upstream of each promoter region (31). The sequence contains a 6-bp inverted repeat typical of sequences recognized by dimeric DNA-binding proteins containing helix-turn-helix motifs. An imperfect inverted repeat upstream of the gsiA promoter region (TTGCGG-N₄-CCGAAA) is homologous to the symmetrical sequence in the degQ and srfA promoter regions (Table 4). Directed mutagenesis studies have revealed that alteration of positions 3 and 4 of the CCGCAA

TABLE 3. Catabolite-resistant sporulation of a gsiA mutant

Medium		CFU	Fre-	
	Relevant genotype	Total	Heat resistant	quency
DS	gsiA ⁺	4.0×10^{8}	2.6×10^{8}	0.65
	gsiA69::cat		2.8×10^{8}	0.72
	AgsiA69::cat SPβgsiA+	3.6×10^8	2.4×10^8	0.67
DS-2% glucose	gsiA ⁺	1.2 × 10 ⁹	1.7 × 10 ⁶	1.4 × 10
	Acri 460cat	8.6×10^{8}	4.9×10^{8}	0.57
	ΔgsiA69::cat SPβgsiA+	1.0×10^{9}	1.2×10^{6}	1.2×10^{-1}

^a Heat-resistant CFU divided by total CFU. The values given are averages of at least two independent experiments.

TABLE 4. Comparison of some ComA-dependent promoter sequences

Gene ^a		Seq	uence		Position ⁶
srfA	TTGCGGC	ATCCCGCAA-I	-TTGCT	PTAAATAAACT	-117
srfA				CCATITITCG	-73
degQ				PACTTTTCGGT	-70
gsiA				PITATITIGG	-75
ComA box	TTGCGGnnnnCCGCAA-Hn-TTGCnnnnATTT				
Conserved frequency	443444	344342	4434	3343	

^a The sequence information was taken from the following sources: degQ, Msadek et al. (25); srfA, Nakano et al. (30).

b Location of the 5' base relative to the start point of transcription.

motif prevent ComA-dependent transcription of srfA (31). These nucleotides are invariant in the putative ComA boxes identified upstream of the srfA, degQ, and gsiA promoter regions (Table 4). This reinforces the notion that this sequence functions as a cis-acting target site for activation by

a ComA-dependent mechanism.

The comP, comQ, spo0K, and comA products appear to function as components of a signal transduction system that results in phosphorylation of ComA by ComP and that in turn causes increased expression of srfA with subsequent activation of late genes in the competence cascade (14, 33, 50). This cannot be the only pathway for phosphorylation of ComA or the only pathway derived from the activity of ComA, however. Transcription of gsiA and degQ (26), while dependent on ComA and ComP, is independent of srfA. Moreover, mutations in degQ or gsiA have no obvious effect on competence development (26). For gsiA, the necessity for positive regulation mediated by ComP-ComA can be efficiently bypassed by a mecA mutation. If the effect of the mecA mutation on gsiA is direct and if the allele of mecA tested has a gain-of-function mutation, such as a mutation that causes constitutive synthesis or activation of MecA, this result might imply either that MecA normally interacts with the gsiA promoter region or that MecA can acquire by mutation the ability to activate gsiA transcription. If the mecA42 allele is a loss-of-function mutation, MecA might normally be a negative regulator of gsiA transcription whose activity must be counteracted by ComA to allow gene expression. Direct interaction between the gsiA promoter and ComA or MecA will be demonstrated only when it is possible to do DNA-binding experiments by using purified proteins. Regulation of the gsiA, com, and deg systems clearly has overlapping elements, although each system has unique features as well. For example, com and deg genes respond to both the ComA-ComP and DegU-DegS systems (albeit in different ways), while gsiA responds to only ComA-ComP. Thus, transcriptional control of adaptive responses in B. subtilis appears to involve coordination of multiple regulatory pathways.

We have little information on the precise function of GsiA; however, in the accompanying report (28) we show that the turnoff of gsiA transcription after the first hour of the stationary phase fails to occur in certain early-blocked sporulation (spo0A and kinA) mutants. The sporulation defect in kinA mutants results from overexpression of gsiA and indicates that 2 product of the gsiA operer is a negative

regulator of an early developmental event.

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